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Organophilic compositions.

There is disclosed an organophilic composition which comprises a mixture of a smectite clay and a
perficulate second inorganic material, the mixture having been treated with a quaternary ammonium compound
which is capable of rendering the smectice lay organophilic and the second inorganic material constituting at
least 10% by weight of the mixture of smectite clay, second inorganic material and quaternary ammonium

The organophilic composition is useful for gelling organic solvents and compositions containing organic solvents.

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ORGANOPHILIC COMPOSITIONS

This invention relates to organophilic compositions and more particularly but not exclusively is concerned with organophilic compositions autiable for improving the thiotropic properties of cross-in-table unsaturated polyester resin compositions and for gelling organic solvents and compositions containing significant amounts of organic solvents. The present invention is also concerned with a process for preparing organophilic compositions.

BACKGROUND OF THE INVENTION

10 Unsaturated polyester resin compositions containing gelling agents, or gellants, are well-known. For example, in the preparation of glass fibre laminates using unsaturated polyester resins as the bonding agent, it is known to use thickropic gelling agents which permit asymmizing and spraying of the resins at high shear rates to which greatly increase the viscosity of the resins at low shear rates thus inhibiting the drainage of the resins on vertical surfaces. One of the most effective golling agents for unsaturated problester resins is a stilice aerogel which is typically added to a solution of the resin in an unsaturated aromatic monomers, such as styrene, at a level of about 1% by weight, based on the weight of the solution. However, slice aerogels are extremely fine and very expensive and, because of their low bulk density, present storage and handling problems and constitute a health hazard.

Various attempts have been made to use organoclays as gelling agents for unsaturated polyester resins, which organoclays are generally smectite clays, e.g., montmorillonite, bentonite, hectorite or the like, at least part of the exchangeable cations of which are replaced by quaternary ammonlum cations containing at least one alkyl group having from 10 to 24 carbon atoms.

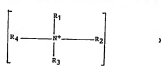
25 SUMMARY OF THE INVENTION

According to one aspect of the present invention, there is provided an organophilic composition which comprises a mixture of a smectite clay and a particulate second inorganic material, the mixture having been treated with a quaternary armnonium compound which is capable of rendering the smectite clay or capacitative and a second inorganic material constituting at least 5% by weight of the mixture of smectite clay, second inorganic material and quaternary armnonium compound.

The amecitic clay may be, for example, bentonite, montmorillorite, hectorite, saponite or fullers earth. The second inorganic material may be, for example, finely divided silica having an average particle claimater in the range from 0.005 um to 0.5 um. Preferably his second inorganic material constitutes from

35 10% to 50% by weight of the mixture of smecifits clay, second inorganic material and customary ammonium compound.

Advantageously, the quaternary ammonium compound has at least one alkyl radical having from 10 to 24 carbon atoms and most preferably is one which can be represented by the general formula:



wherein R, is a saturated or unsaturated alkyl group having from 10 to 24 carbon atoms, R, and R, which may be the same or different, are each a saturated or unsaturated alkyl group having from 1 to 24 carbon atoms or an arakyl group having from 7 to 10 carbon atoms, R, is an alkyl group having from 1 to 8 carbon atoms or an arakyl group having from 7 to 10 carbon atoms, and X is OH, Cl, Br, I, No., CH,SO, or CH,cOO. Examples of such quaternary ammonium compounds are the metryl benzyl disklyl ammonium

chlorides, the dimethyl disklyf ammonium chlorides, the dimethyl benzyl alkyl ammonium chlorides, the benzyl triaklyl ammonium chlorides and the methyl triaklyl ammonium chlorides in which the alkyl group is a mixture of hydrocurbon radicals derived from tallow having from 14 to 20 carbon atoms but in which C_n radicals predominate. (A hypical analysis of such a mixture of hydrocarbon radicals contained in tallow is: 5 C_n 4.5%; C_n 0.5%; C_n 0.5%; C_n 1.5%; C_n 6.20% and C_n 1.0%.) The hydrocarbon radicals may be partially unsaturated as in natural tallow or may be substantially completely saturated as a result of treating the tallow with hydrogen in the presence of a suitable catalyst:

Most preferably, the mixture of smectite clay and second inorganic material is treated with a mixture of distributed difflydrogenated tallow ammonium chloride (2M2HT) and dimethyl benzyl hydrogenated tallow ammonium chloride (2M3HT) in proportions ranging from 25 mol % of 2MBHT and 75 mol % of 2M2HT to 100 mol % of 2M2H and 0 mol % of 2M2HT to

It is also preferred to treat the mixture of smectits clay and second inorganic material with a quantity of quaternary ammonium compound or mixture of quaternary ammonium compounds such as to provide from 95 to 140 milliequivalents of quaternary ammonium compound(s) per 100 g of dry smectitle clay.

7.5 According to another aspect of the present invention there is provided a process for preparing en organophilic composition, which process comprises preparing an aqueous suspension comprising a mixture of a smectite clay and a particulate second inorganic material, mixing with this aqueous suspension a questionary ammonium compound capable of rendering the smectite clay organophilic, either in the motion state or in the form of an emulsion in water, and subjecting the resultant mixture to high shear mixing for a second inorganic material being used in an amount sufficient to provide at least 5% by weight of the final mixture of smectite clay, second inorganic material abort questions.

In the process of the present invention, the high shear mixing is preferably effected by passing the suspension through a homogenizer of the type in which the suspension is forced in the form of a thin film segments through a thin, hard surfaced gap under a pressure of at least 250 pounds per square inch (1.7 for in United States Patent Specification Nos. 397/76 for in United States Patent Specification Nos. 3037/73 by the Manton-Gaulin Company. Advantageously, the homogeniser is pressure of at least 250 pounds per square inch (10.5 MPa). The amount of energy, in KJ per Kg of dry solids, dissipated in the mixture is given by the expression:

$$E = nP.10^3$$

where P is the pressure in MPa exerted in the Manton-Gaulin homogeniser, a is the number of passes through the Manton-Gaulin homogeniser, and w is the weight in grams of dry solids in 1 litre of the aqueous moture.

The mixture of smecitie clay and second inorganic material after reaction with the quaternary ammonium compound(s) is preferably dewatered by filtration or by centrifugation, washed with water and thermally dried under conditions such that the temperature does not exceed 100°C for more than a term minutes. The dry material may then be pulverised to facilitate its incorporation into, for example, a cross-include unsaturated polyester resin composition. Alternatively, the minuter of smectite clay and second inorganic material after reaction with the quaternary ammonium compound(s) may be pertially dewatered for, example, by decantation and the hickness suspension thus formed spray dried to give dry material in the form of microspheres which may not require any pulversation.

The organophalic composition of the invention is also suitable for gelling organic solvents such as tousne and sylene and compositions containing such solvents, and for trickening polar paint compositions such as "two-pack spoxy paint". Polar paint compositions are generally palints which contain organic solvents of medium to high polarity, such as ketones, esters, glycol ether esters and alcohols. It is important that the organocity thickener should disperse quickly and easily into the peint composition with the consumption of the minimum amount of energy. This is reactly achieved using the organophilic composition of the present invention.

EMBODIMENTS OF THE INVENTION

The invention is illustrated by the following Examples.

EXAMPLE 1

An aqueous suspension of bentonite was prepared by mixing raw Wyoming sodium bentonite in a blunger with sufficient water to form a suspension containing 101% by weight of dry clay and with 1% by weight, based on the weight of dry clay, of terzeodium pryphosphate as a dispersing agent. The deflocutated suspension thus formed was passed through a No. 300 mesh British Standard slove (nominal aperture 0.055mm) and the undersize fraction was diluted to about 6% by weight of solids with water and subjected to a particle size separation in a nozzle discharge, disc centrifuge at a flow rate of 2 litres per minute. The fine fraction was then passed once through a Mantion-Qualin homogeniser at a pressure of 4000 psi (27.6 MPa). The energy dissipated in the suspension during the single pass through the Mantion-Gaulin homogeniser was 673 kJ per Kg of dry clay.

Three aqueous suspensions A. B and C were prepared each containing 6% by weight of a precipitated silica having a mediate utilimate particle diameter of 0.03um. All three suspensions were formed initially by mixing the silica with the water by means of a laboratory stirror.

Suspension A was then further treated by being passed once through the Manton-Qaulin homogeniser at a pressure of 4000 psi (27.6 MPa), the energy dissipated in the suspension being 443 KJ per Kg of dry cilica.

Suspension B was further treated by means of a Silverson shrouded impeller mixer set to provide maximum shear for a time of 15 minutes. The energy dissipated in the auspension under these conditions as leavey difficult to estimate but it is believed to be of the order of 100 KLJ per KJ.

Suspension C was given no further mechanical treatment.

The bentonite suspension was indeed with a suspension of silica in proportions such that the mixture contained 25% by weight of dry silica in the final dry organoclay/silica product and 48.9% by weight of dry clay. The remaining 28.1% is accounted for by the dry weight of the mixture of quaternary ammonium of the suspension of a mixture consisting of 50 mol % of dimetrity diffly/dropensted tallow) ammonium chloride (2MBCHT) and 50 mol % of dimetrity benzy (invitropension diffly/surpopension different diffly/surpopension different diffly/surpopension different diffly/surpopension different different

Mbdures of bentonite suspension, silica suspension and quaternary ammonium compounds were prepared by the following methods:

- The bentonite and silica suspensions were mixed together in the desired proportions using the Silverson mixer at the maximum shear setting for 15 minutes, during which time sufficient of the emulation of mixed quaternary ammonium compounds was added to provide 105 milliequivalents of queternary ammonium compounds per 100g of dry bentonite.
- 2. The bentonite and silica suspensions were mixed together using the Manton-Gaulin homogenizer for one pass at 4000 psi (27.5 MPa) and the emulsion of mixed quaternary ammonium compounds was added to the feed container of the Manton-Gaulin homogenizer in the same proportions as in method 1. The energy dissipated in the suspension was 1856 KJ per Kg of total dry solids.
 - 3. The bentonite and silica suspansions were mixed together using the laboratory stimer and the emulsion of mixed quaternary ammonitum compounds was added as the stirring proceeded in the same proportions as in metiod 1.
- 4. The bentonite suspension and the emulsion of mixed quaternary ammonium compounds were mixed together using the Manton-Gaulin homogeniser for one pass at 4000 psi (27.6 MPa) and the suspension or sitica was added to the feed container of the Manton-Gaulin homogeniser. The energy dissipated in the suspension was 1024 KJ per Kg of total dry solids.

A further mixture was prepared, as a control, by mixing together the bentonite suspension only and sufficient of the emulsion of mixed quaternery emmonium compounds in proportions such as to provide 105 si milliequivalents of quaternery emmonium compounds per 100g of dry bentonic testing the Manton-Gaulin homogeniser for one pass at 4000 psi (27.6 MPa). The energy dissipated in the suspension was 1402 kJ per Kg of total dry solids.

in each case the organically suspension was filtered on a Buchner funnel, washed with hot water and dried for 16 hours at 60 °C in an air-swept oven. The dry product was then milled to pass through a seive of nominal aperture 0.080mm.

1% by weight of each dry organoclay composition was incorporated into an unsaturated polyester resin 5 composition using a laboratory stirrer rotating at 3,500 rpm. The resin was a rapid-curing, general—purpose polyester resin marketed by the Scott Bader Company Limited under the trade name "CRYSTIC 196". ("CRYSTIC" is a Registered Trade Mark), and, is believed to be a co-polymer of g-phthalic acid, fumaric acid and propylene glycol, having a number average molecular weight of about 3000.

The viscosity of the polyester resin composition containing the organoclay as a gelling agent was 10 measured using a Brookfield Viscometer at a spindle speed of 0.5 rpm.

The results obtained are set forth in Table I.

15		TABI	TABLE I		
ra	Silica suspension	Mixing <u>Method</u>	Viscosity (mPa.s	Thixotropy Index	
20	No silica	Manton- Gaulin	1300	2.30	
	C	3	600	1.80	
	В	1	2800	2.78	
25	A	2	3200	2.78	
~	A	3	800	2.00	
	λ	4	1280	2.30	

These results show that in order to achieve a good galling effect in the polyester resin, the silica suspension must be mixed with the bentonite suspension before the bentonite is contacted with the quaternary ammonium compounds, and that both the silica suspension and the bentonite/silica/quaternary ammonium compound mixture must be subjected to high shear mechanical working such that at least 100 KJ of energy per Kg of dry total solids is dissipated in the suspension.

The thixotropy index referred to in Table 1 is defined as the ratio of the viscosity of the composition as measured with a Brookfield Viscometer at 5 rpm to the viscosity measured with a Brookfield Viscometer at 50 rpm. The value of the thixotropy index should be as high as possible where shear thinning or pseudoplastic properties are sought.

EXAMPLE 2

The bentonite suspension and silica suspension "A" prepared as described in Example 1 above were mixed together in proportions such as to give 25% by weight of dry silica in the final dry organoclasy/silica product and in the range of from 46.3% to 51.8% by weight of dry bentonite, depending upon the effective molecular weight of the mixture of quaternary ammonium compounds, and samples of the mixed suspension were passed once through the Manton-Gaulin homogeniser at 4000 psi (27.6 MPa). In each case there was added to the feed container of the Manton-Gaulin homogeniser sufficient of an emulsion containing 1% by weight of a mixture of 2M2HT and 2MBHT to provide 105 milliequivalents of mixed quaternary ammonium compounds per 100g of dry bentonite, but the 2M2HT and the 2MBHT were mixed in different

in each case the organoclay suspension was filtered, washed, dried and milled as described in Example 1 and 1% by weight of each dry organociay was stirred into the same polyester resin composition as was described in Example 1. The viscosity of each polyester resin sample after the addition of the organically se was measured by means of a Brookfield Viscometer at a spindle speed of 0.5 rpm and the results obtained

TABLE II

	mole % of		% by weight of				
5	2MBH1	r 2M2HT	bento- nite	quarter- nary ammonium compounds	(mPa.s)	Thixotropy Index	
10	0	100	46.3	28.7	1200	1.90	_
	25	75	47.5	27.5	1800	2.40	
	50	50	48.9	26.1	3200	2.78	
15	75	25	50.3	24.7	3800	2.9	
	100	0	51.8	23.2	3600	2.8	

- The bentonite suspension and silics suspension "A" prepared as described in Example 1 above were mixed bigether in proportions such as to give 28% by weight of dry silics and 48.9% by weight of dry bentonitie in the final dry organocisystilics product, the remaining 28.1% being accounted for by the dry weight of the mixture of quaternary ammonium compound. Samples of the mixed suspension were passed once through the Manton-Gaulin homogeniser at 400 psi (27.6 MPz), in each case there was added to the feed container of the Manton-Gaulin homogeniser a quantity of an emulsion containing 1% by weight of a mixture consisting of 75 md ½ of 2MBHT and 25 mm ½ or 2BMT. The quantity being varied in each case in order to provide a different number of milliequivalents of mixed quaternary ammonium compounds per 100 g of dry bentonite.
- in each case the organoclay suspension was filtered, washed, dried and milled as described in Example 1, and 1% by weight of each dry organoclay was stirred into the same polyester resin composition as was described in Example 1. The viscosity of each polyester resin after the addition of the organoclay was measured by means of a Brookfield Viscometer at spindle speed of 0.5 rpm and the results obtained are set forth in Table III.

		T	ABLE III		
	Milli- equivalents	% by we	% by weight of		
40	of quarter- nary ammonium compounds per 100g of bentonite	bento- nite	quarter- nary ammonium compounds	Viscosity (mPa.s)	Thixotropy Index
45	85	53.7	21.3	300	1.15
	90	52.8	22.2	800	1.90
	95	51.9	23.1	2300	2.50
50	100	51.1	23.9	3200	2.80
	105	50.3	24.7	3800	2.90
	110	49.5	25.5	3400	3.10
55	115	48.8	26.2	3600	3.00
	120	48.0	27.0	3000	2.50

The bentonite suspension and silica suspension "B" prepared as described in Example 1 above were mixed together in various proportions to give mixed suspensions each containing a different percentage by weight of silica. Each mixed suspension was prepared using the Silverson mixer at the maximum shear setting for 15 minutes. In each case there was added to the mixed suspension as mixing proceeded a quantity of an emulsion containing 1% by weight of a mixture consisting of 50 mol f. of 2MBHT and 50 mol 10 % of 2MBHT sufficient to provide 105 milliequivalents of mixed quaternary ammonium compounds per 100g of dry bentonite.

In each case the organoctay suspension was filtered, washed, dried and milled as described in Example 1, and 1% by weight of each dry organoctay was stirred into the same polyester resin composition as was described in Example 1. The viscosity of each polyester resin after the addition of the organoctay was remained by means of a Brookfield Viscometer at a spindle speed of 0.5 rpm and the results obtained are set tooth in Table IV.

TABLE IV
Percentage by weight of

25	silica	bentonite	quarternary ammonium compounds	Viscosity (mPa.s)	Thixotropy Index
30	0 10 25 40 75 100	65 58.7 48.9 39.0 16.3	35 31.2 26.1 21.0 8.7	2000 2500 2800 2000 400 400	2.35 2.57 2.78 2.50 1.25 1.02

EXAMPLE 5

Two-pack epoxy enamel paint

An aqueous suspension of bentonite was prepared by mixing raw Wyoming sodium bentonite in a blunger with sufficient water to form a suspension containing 10% by weight of dry clay. The suspension flus formed was feed to a scroll-type centrituge which removed substantially all of the particles targer than 50 microns. The degritted bentonite suspension was then diluted with water to a solids content of 4% by weight and passed through a No. 300 mesh British Standard sieve. The undersize fraction was subjected to a particle size separation in a nozled discharge, disc centrituge at a flow rate of 1 litre per minute. The fine fraction produced by the cnetrifuge had a solids content of 2.5% by weight.

An aqueous suspension of precipitated silica was also prepared by mixing with water by means of laboratory stime sufficient of a precipitated silica having an average particle diameter of 0.4 micron and a specific surface area of 200m⁴0" to form a suspension containing 4.3% by weight of dry silica.

The bentonite fine fraction and the precipitated silica suspension were mixed together in proportions such that the final dry organockay product contained 20% by weight of silica and the mixed suspension was passed once brough the Manton-Gaulin homogenisor at a pressure of 4000psi (27.6 MPa), the energy dissipated in the emulsion being 988 KJ per kg of dry solids.

An aqueous supposton of 2MBHT was prepared by melting the quaternary ammonium compound/sopropyl alcohol mixture which is available in commerce and pouring the molten mixture into hot water in proportions such as to form a 1% by weight remulsion of 2MBHT in water. The emutation was passed once through the Manton-Gaulin homogeniser at a pressure of 4000 psi (27.6 MPa), the energy dissipated in the emulsion being 279s KJ per kg of dry softly.

Portions of the suspension of the bentonite/silica mixture were mixed with portions of the emulsion of 2MBHT emulsion at 65°C in a paddle mixer to give the proportions of silica, bentonite and quaternary ammonium compound given in Table V, and in each case the mixture was agitated for 30 mixtures and then passed once through the Manton-Gaulin homogeniser at a pressure of 4000 psi (27.6 MPA). Each or organociary portion was then filtered, dried in an air swept oven for 16 hours at 60°C and milled to pass through a sieve of nominal aperture 0.80 mixed.

Each portion was incorporated into pack A of a two pack white epoxy paint, of which pack A is the base containing the epoxy resin and pack B is the polyamide curing agent solution.

Pack A was prepared by mixing together the following ingredients which are listed in their order of addition:

		Grams
5	75% by weight solution of epoxy resin	
	in xylene	157.0
	Solvent mixture (see below) Urea formaldehyde resin as flow	47.5
	control agent	7.0
,	Dispersing agent	3.5
	Titanium dioxide pigment	149.0

The above ingredients were mixed together in a water-cooled pot using a 4 cm diameter Cowles blade at 4000 rpm. The mixing was continued until a small sample spread on a Hegman gauge gave a reading of less than 5 micross.

The speed of the Cowles blade was then reduced to 2000 rpm and there were then added:

i		Grans
	Solvent mixture (see below)	42.5
	Organoclay	2.5

When all the organocial had been added the speed of the Cowles blade was again increased to 4000 pm and, for each batch of pack A, small samples were taken at 1 minute intervals and spread on a hegman gauge. For each batch there was recorded the time taken to reach a state of dispersion such that the Hegman gauge gave a reading of less than 10 microns (see Table V below).

As a comparison, further batches of pack A were prepared containing organoclays prepared in a manner similar to that described above but containing no sitica and the mixing time required to reach a Hegman gauge reading of less than 10 microns was recorded (see Table V below).

Each batch of pack A was mixed with a batch of pack B prepared according to the following formulation, to form a white epoxy enamel paint

				Grams
Polyamide Xylene	liquid	curing	agent	56.0 35.0

The total formulation weight of the epoxy paint was 500 grams. The solvent mixture comprised:-

	Ethylene glycol monoethyl	* by weight
,	ethyl acetate	65
	Methyl isobuty ketone	15
	Xylene	20

50

The results obtained are set forth in Table V.

anneu a	7 mm to reach place to reach place to reach place to reach place that the search of th
	Energy dissipated in homogeniser (KJ per ed ary 1371. 1371. 1396. 1434. 1549. 1552. 1652. 1652.
TABLE V	#by weight of bentonike quarternary emmoatum compound compound 57.6 22.4 53.8 26.1 23.9 52.0 28.0 72.5 \$7.5 \$7.5 \$7.5 \$7.5 \$7.5 \$7.5 \$7.5 \$7
	meq of quarternary menonium compound per 100g of bentonite 91 106 116 89 101 1113
! ^ ! !	######################################

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EXAMPLE 6 Galling toluene

5 Aqueous suspensions of bentonite and precipitated silica were prepared using the same starting materials and preparative methods as described in Example 5.

The suspension of finely divided bentonite and the suspension of precipitated silica were mixed together in different proportions and each mixed suspension was passed once through the Manton-Gaulin homogeniters at a pressure of 276 MPa.

An aqueous suspension containing 2MBHT and 2M2HT in the proportions 75 mol % 2MBHT to 25 mole % 2M2HT was prepared by melting the quaternary ammonium compound/sopropyl alcohol mixtures and pouning the molten mixture into hot water to form an emulsion containing 1% by weight of the mixed quaternary ammonium compounds in water.

Portions of the amulsion of mixed quaternary ammonium compounds were circulated through the Manton-Gaulin homogeniser at a pressure of 27.8 MPz and in reach case the bentiontal/precipitated silice suppension was action to the feed hopper of the homogeniser in proportions such that there were present in the mixture 105 med of quaternary ammonium compounds/100 g of dry bentionits, in each case the mixture received the equivalent of one pess through the homogeniser. The organoclasy/silice composition to the formed was then filtered, washed, dried in an air-swept oven at 80 °C for 18 hours and milled to pass through a selver of nominal abenture 0.080 mm.

Each sample of dry organoclay/allica composition was tested for its gelling properties in toluene by dispersing a 6 g sample of the composition in 340 ml of toluene in a Waring Blendor rotating at 16500 pm for 3 minutas. The mixture was allowed to stand for 2 hours and the viscosity than measured using a Brockfield Viscometer at a speed of 50rpm. The results obtained are set out in Table VI

65

		(mpas) 200 210 210 200 290 330 450 450 520
ated	r organoclay/	111ca 1639 1589 1587 1487 1386 1335
Energy dissip	in homogeniser (KJ Kg) in silica/	1069 1069 1055 1053 992 963 908 882
clay/silica	quaternary ammonium compounds	32.9 31.3 29.6 28.0 26.3 24.7 23.1
by Weight in organoclay/silica omposition of	bentonite	63.7 63.7 60.4 57.0 53.7 50.3 46.9
omposition of	ilica	

TABLE VI

25

Aqueous suspensions of bentonite and precipitated silica were prepared as described in Example 5 and were mixed together in proportions such that the final organoclay/silica composition contained 20% by seeight of dry silica. Each mixed suspension was passed once through the Manton-Gaulin homogeniser at a pressure of 27.6 MPa.

A 1% by weight aqueous emulsion of a mixture consisting of 75 mole % 2MBHT and 25 mole % 2M2HT was prepared as described in Example 8. Portions of this emulsion were circulated through the Manton-Gailla homogeniser at a pressure of 27.8 MPa and in each case a bentonite/precipitated silica to suspension was added to the feed hopper of the homogeniser in proportions such that in each case there was present a different number of milliequivalents of quaternary ammonium compounder100 g of dry bentonite. In each case the mixture received the oquivalent of one pass through the homogeniser. The for 16 hours and millied to pass through a there were then filtered, washed, dried in an air-swept oven at 60°C for 16 hours and millied to pass through a time serve of norminal sperture 0.000 mm.

Each sample of dry organoclay/sitica composition was tested for its gelling properties in toluene by the method described in Example 6 and the results obtained are set forth in Table VII below:

KΛ

TABLE VII

J Viscosity 1V/(mPas)	800 530 350 124 40
Energy dissipitated in homogeniser (KU kg-1 in Viscos silica/ organoclay/(mPas)	Dentonite silica 967 1392 964 1424 963 1447 959 1476
Energy d in homog (KJ kg-1 silica/	967 963 963 962 959
<pre>% by weight in organoclay/ silica composition of benconite quaternary ammonium componing</pre>	23.7 25.6 26.3 27.0 28.7
	56.3 54.4 53.7 53.0 51.3
meg quaternary ammonium compounds Per 100 g bentonite	90 100 110 120

20

Aqueous suspensions each consisting of a mixture of water bentonite and precipitated silica were prepared as described in Example 7. Each mixed suspension contained bentonite and silica in proportions such that the final organochyelica composition prepared from each contained 25% by weight of dry silica and was passed once through the Manton-Gaulin homogeniser at a pressure of 27.6 MPa.

Emulsions were prepared containing 1% by weight of mixtures in various proportions of 2MBHT and 2M2HT according to the method described in Example 6. Each batch of emulsion was circulated through the Martion-Gaulin homogeniser at a pressure of 27.6 MPa and in each case a bentonite/precipitated silica to suspension was added to the feed hopper of the homogeniser in proportions such that there were present 105 meq of quatemery ammonium compounds per 100 g of dry bentonite. In each case the mixture received the equivalent of one pass through the homogeniser. The organoclay/silica compositions thus formed were their filtered, washed, dried in an air-swept oven at 80°C for 16 hours and milled to pass through a sleve of nominal sperture 0.800 mm.

Each sample of dry organoclay/sitics composition was tested for its getting properties in toluene by the method described in Example 8 and the results obtained are set forth in Table VIII below:

Viscosity (mpas)	430 190 30 30
Energy dissipitated in homogeniser (KJ kg-1) in silical	Dentonite silica 938 1361 932 1409 929 1409 926 1453
* by weight in organoclay/ silica composition of bentonite ammonium commonium	51.8 50.3 23.2 48.5 26.1 47.5 27.5 46.3
Mole % in quaternary ammonium compound mixture of ZMBHT ZMZHT	100 0 75 25 50 50 25 75 0 100

Claims

- 1. An organophilic composition, characterised in that it comprises a mixture of a smecitie clay and a particulate second inorpanic material, and in that the mixture has been treated with a quaternary ammonium compound which is capable of rendering the smecitie clay organophilic and the second inorpanic material constituting at least 5% by weight of the mixture of smecitie clay, second inorganic material and quaternary ammonium compound.
- A composition as claimed in claim 1, wherein the second inorganic material is finely divided silica having an average particle diameter in the range from 0.005 µm to 0.5 µm.
- 3. A composition as claimed in claim 1 or 2, wherein the second inorganic material constitutes from 10% to 50% by weight of the mixture of smectite cley, second inorganic material and quaternary ammonium compound.
- A composition as claimed in claim 1, 2 or 3, wherein the quaternary ammonium compound has at least one alkyl radical having from 10 to 24 carbon atoms.
- 5. A composition as claimed in claim 4, wherein the quaternary ammonium compound is one which can be represented by the general formula:



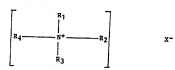
- wherein R, is a saturated or unsaturated alkyl group having from 10 to 24 carbon atoms, R, and R, which may be the same or different, are each a saturated or unsaturated alkyl group having from 1 to 24 carbon atoms or an arakly group having from 7 to 10 carbon atoms, R, is an alkyl group having from 1 to 8 carbon atoms or an araklyl group having from 7 to 10 carbon atoms, and X is OH, Ci, Br, I, NO, Ch,SO, or CH,SOO.
- 5 6. A composition according to claim 1, 2, 3, 4 or 5, wherein the mixture of smecitie clay and second inorganic material is reased with a mixture of dimethyl diffrydrogenated tailow) ammonium chloride. and dimethyl benzyl hydrogenated tallow ammonium chloride.
- 7. A composition according to claim 6, wherein said mixture comprises from 25 mol % to 100 mol % of dimethyl benzyl hydrogenated tellow ammonium chloride and from 75 mol % to 0% of dimethyl di-(fhydrogenated tallow) ammonium chloride.
 - 8. A composition as claimed in any one of claims 1 to 7, wherein the mixture of smecitie clay and second inorganic material is treated with a quantity of quaternary ammonium compound or mixture of quaternary ammonium compounds such as to provide from 95 to 120 milliequivalents of quaternary ammonium compound(s) per 100g of dry of smecitie day.
- 9. A process for preparing an organophilic composition, characterised in that the process comprises preparing an equeues suspension comprising a mixture of a smectite city and a particulate second inorganic material, mixing with the acqueous suspension a questernary ammonium compound which is capable of rendering the smectite city organophilic, either in the molten state or in the form of an emulsion in water, and subjecting the resultant mixture to high shear mixture, and in that the second inorganic material is used in an amount sufficient to provide at least 5% by weight of the final mixture of smectite city, second longrapic material and questernary ammonium compound.
- 10. A process according to claim 8, wherein the high shear mixing is effected by passing the suspension through a homogeniser of the type in which the suspen sion is forced in the form of a thin film edgewise through a thin, hard surfaced gap under a pressure of at least 250 pounds per square inch (1.7 MPa) and at a high velocity.

- 11. A process according to claim 9 or 10, wherein after the mixture of smectite clay and second inorganic material has been treated with the quaternary ammonium compound(s) it is devatered by filtration or by centrifugation, washed with water and themally dried at a semperature not exceeding 100°C.
- 12. A cross-linkable unsaturated polyester resin composition including a minor amount of an organophilic composition as claimed in any one of claims 1 to 8.
- 13. A composition comprising an organic solvent and an amount of an organophilic composition as claimed in any of Claims 1 to 8, sufficient to get the organic solvent.
- 14. An epoxy resin base including, as a thickener, an organophilic composition as claimed in any of Claims 1 to 8.

Claims for the following contracting states: AT, ES:

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- 1. A process for preparing an organophilic composition, characterised in that the process comprises mixing a smectite city and a particulate second inorganic material, and treating the mixture with a requesterary ammonium compound which is capable of rendering the smectite clay organophilic and in that the second inorganic material constitutes at least 5% by weight of the mixture of smectite clay, second inorganic material and quaternary ammonium compound.
 - 2. A process as claimed in claim 1, wherein the second inorganic material is finely divided stitica having an average particle diameter in the range from 0.005 µm to 0.5 µm.
- 3. A process as claimed in claim 1 or 2, wherein the second inorganic material constitutes from 10% to 50% by weight of the mixture of smecifie clay, second inorganic material and quaternary ammonium
- A process as claimed in claim 1, 2 or 3, wherein the quaternary ammonium compound has at least one alkyl radical having from 10 to 24 carbon atoms.
- 5. A process as claimed in claim 4, wherein the quaternary ammonium compound is one which can be represented by the general formula:



wherein R. is a saturated or unsaturated alloyl group having from 10 to 24 carbon atoms, R, and R, which may be the same or different, are each a saturated or unsaturated alloyl group having from 1 to 24 carbon atoms or an analoyl group having from 7 to 10 carbon atoms, B, is an alloyl group having from 8 to 10 carbon atoms or an aralloyl group having from 7 to 10 carbon atoms, and X is OH, CI, Br. I, NO, CH,SQ, or CH,COO.

- 6. A process according to claim 1, 2, 3, 4 or 5, wherein the mixture of smectite clay and second inorganic material is treated with a mixture of dimethyl di(hydrogenated tallow) ammonium chloride and dimethyl benzyl hydrogenated tallow ammonium chloride.
 - 7. A process according to claim 6, wherein said mixture comprises from 25 mol % to 100 mol % of dimethyl bearcyl hydrogenated tallow ammonium chloride and from 75 mol % to 0% of dimethyl di-(hydrogenated labor) ammonium chloride.
- 8. A process as claimed in any one of claims 1 to 7, wherein the mixture of ameditie clay and second inorganic material is treated with a quantity of quaternary ammonium compound or mixture of quaternary ammonium compounds such as to provide from 55 to 120 milliequivalents of quaternary ammonium compound(s) per 100g of dry of ameditte clay.
- 9. A process for preparing an organophilic composition characterised in that the process comprises preparing an aqueous suspension comprising a mixture of a smectite clay and a particulate second inorganic material, mixing with the aqueous suspension a quaternary ammonium compound which is capable of rendering the smectite clay organophilic, either in the molten state or in the form of an emulsion in water, and subjecting the resultant mixture to high shear mixing for a time sufficient to dissipate in the

mbture at least 100KJ of energy per Kg of dry solids in the mixture, and in that the second inorganic material is used in an amount sufficient to provide at least 5% by weight of the final mixture of smectite clay, second inorganic material and quastrampy ammonism compound.

- 10. A process according to claim 8, wherein the high shear mixing is effected by passing the s suspension through a homogeniser of the type in which the suspension is forced in the form of a bin film edgewise through a thin, hard surfaced gap under a pressure of at least 250 pounds per square inch (1.7 MPa) and at a high velocity.
- 11. A process according to claim 8 or 10, wherein after the mixture of smectite clay and second inorganic material has been treated with the quaternary ammonium compound(s) it is dewatered by literation or by contribugation, washed with water and thermally dried at a temporature not exceeding 100°C.
 - 12. A process for preparing a cross-linkable unsaturated polyester resion composition, which process coupliness including in the resin a minor amount of an organophilic composition prepared by a process as claimed in any one of claims 1 to 8.
- 13. A process for preparing a gelled organic solvent which process comprises combining an organic solvent with an amount of an organophilic composition sufficient to gel the organic solvent, said organophilic composition being propered by a process as claimed in any of Calime 1 to 8.
 - 14. A process for preparing a thickened epoxy resin base which process comprises combining an organophilic composition prepared by a process as claimed in any one of Claims 1 to 8 with an epoxy resin base.